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Electrochemical Oxidation of PAHs in Water from Harbor Sediment Purification



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Introduction

PAHs (Polycyclic Aromatic Hydrocarbons)

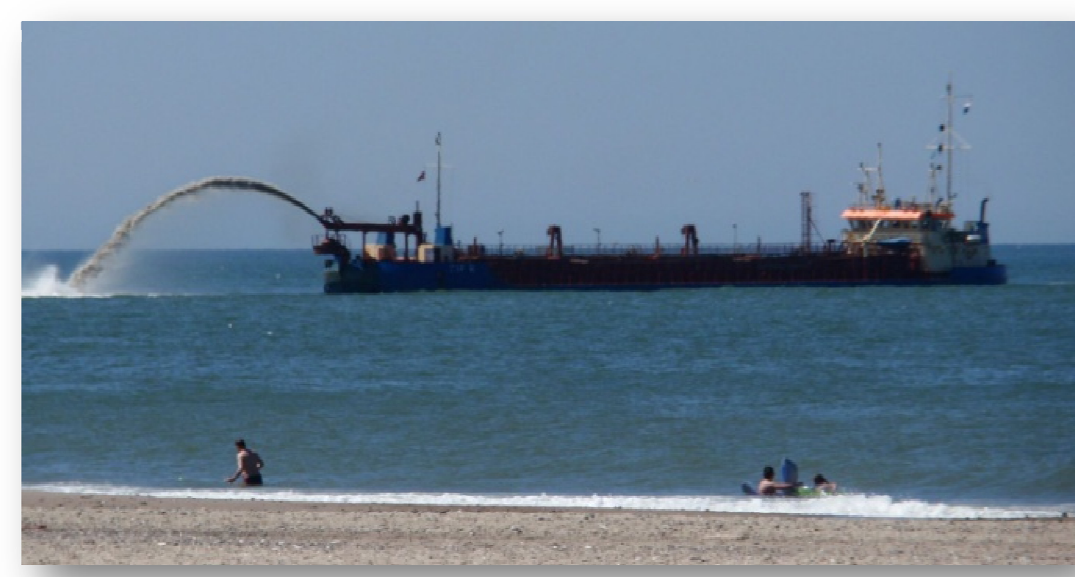
- One of the main POPs (Persistent Organic Pollutants), main sources are burning of fossil fuels (oil and coal), wood stove heating, cigarette smoke, production and use of tar
- Highly toxic, carcinogenic and mutagenic for human beings
- Low threshold values in drinking water: 0.005 - 0.010 µg L⁻¹
- Chemical characteristics: Lipophilic and hence low water solubility, primarily found in soil and sediment, persistent, recalcitrant, and non-reactive

Problem

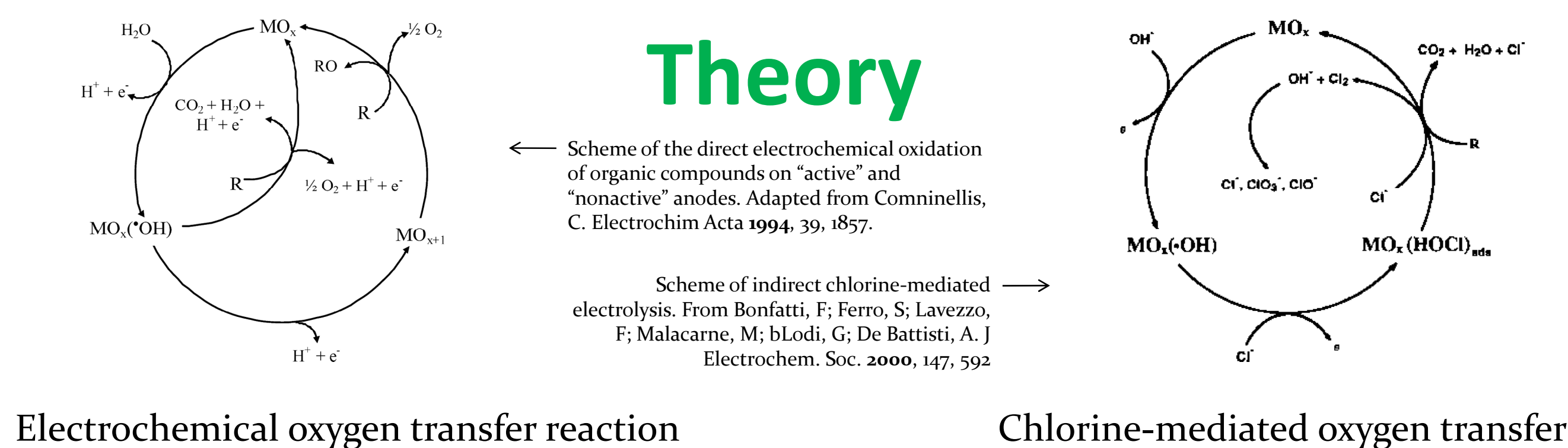
Removal of sediment from harbors produce PAH containing process water, which needs treatment before discharge to the marine recipient.

Objective

Degradation of PAHs in saline process water by means of electrochemical oxidation.

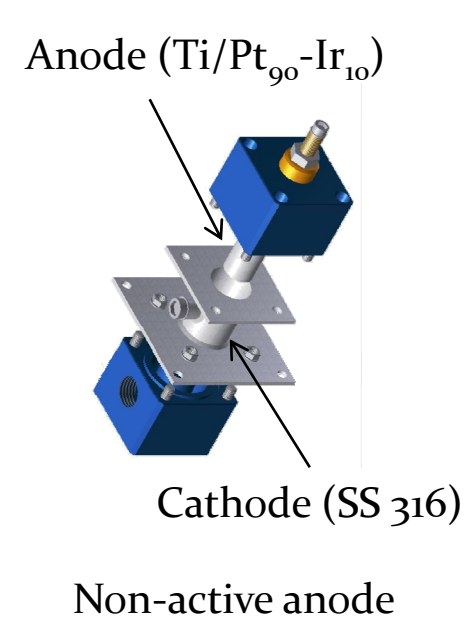


Theory

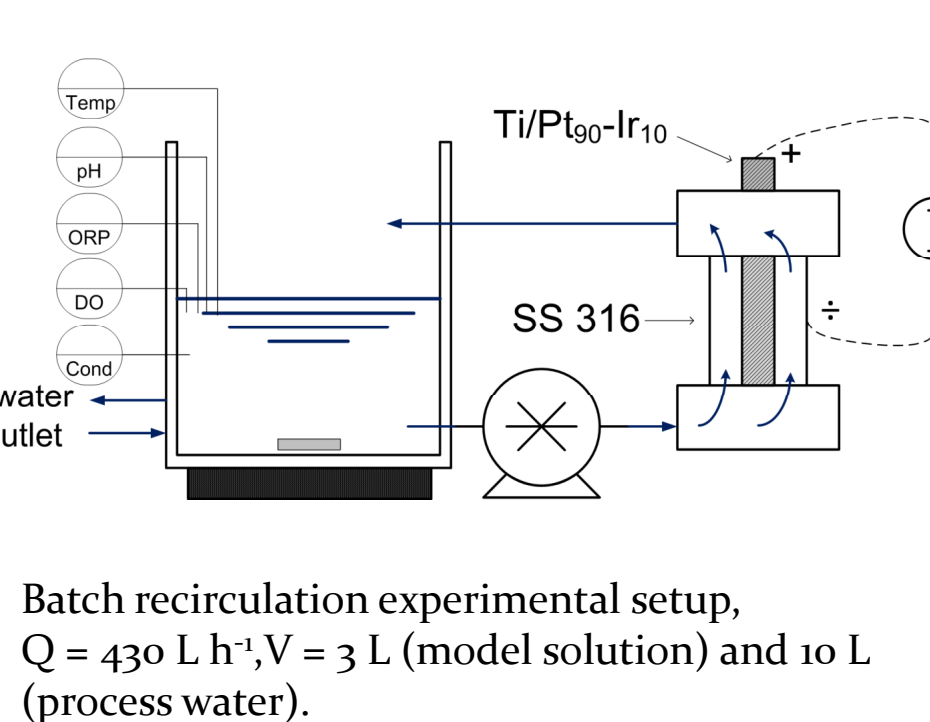


Materials & Methods

The cell



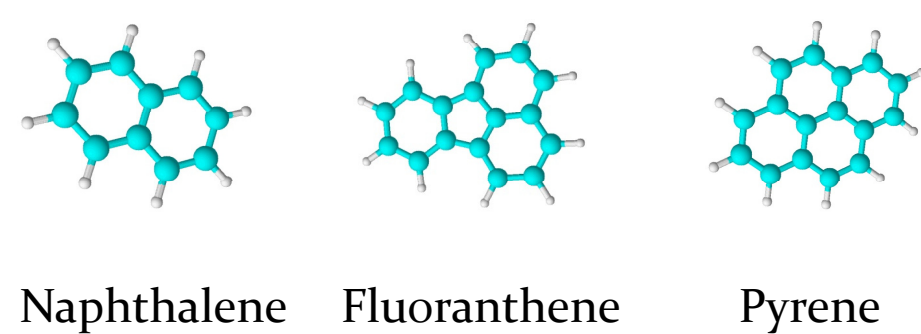
The setup



Process water

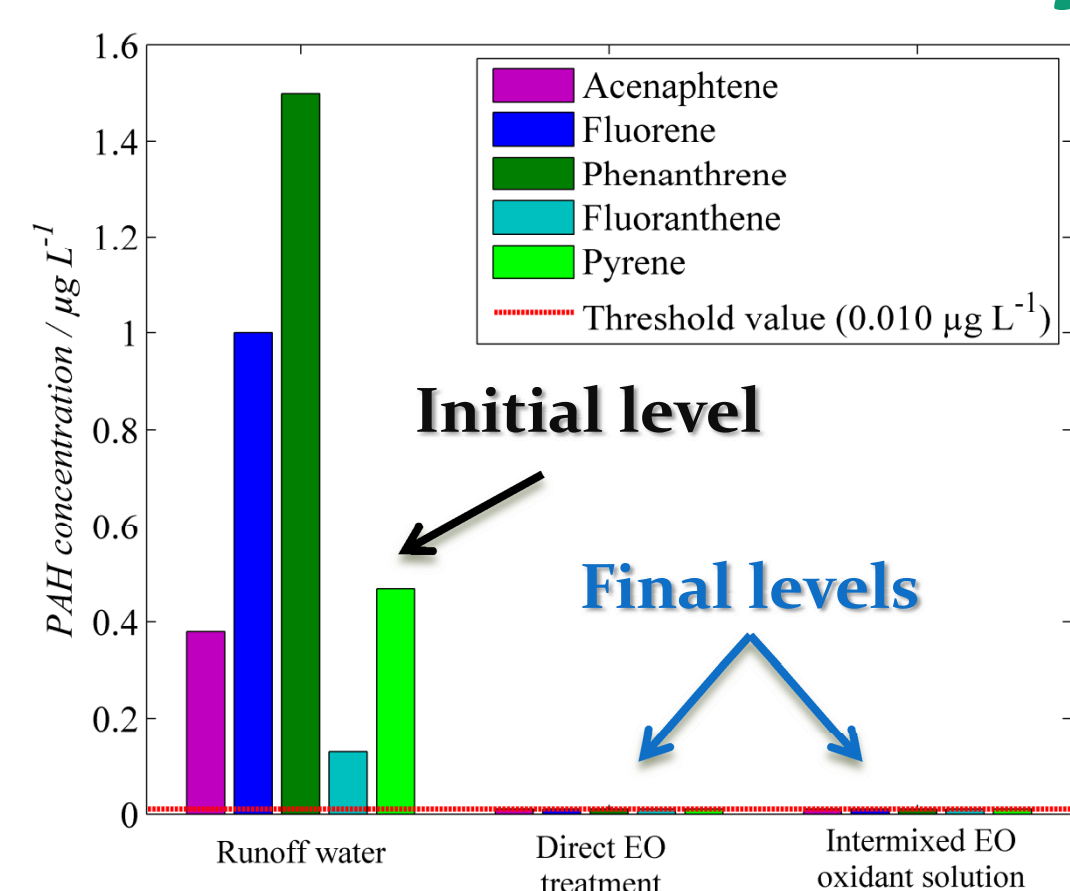
salinity 1 w/w %

Model compounds



Results & Discussion I

Tests on sediment runoff water (process water)



Electrochemical treatment of runoff water from disposal and purification of harbor sediment, which is pumped ashore to upland sites. The initial concentrations of PAHs are low, but treatment is still required to maintain the discharge limits.

Electrochemical treatment approach:

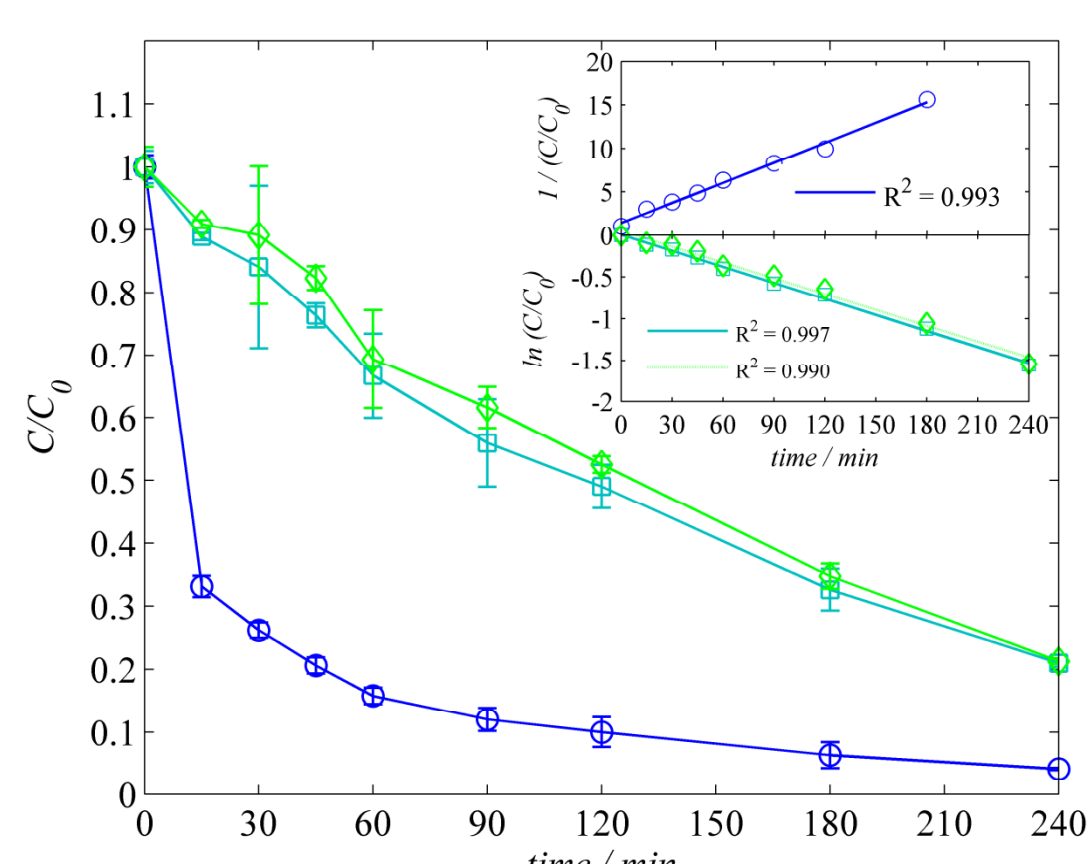
- PAH removal obtained to below discharge limit (0.010 µg L⁻¹)
- Energy consumption: 13.2 kWh m⁻³

Treatment by EO generated chemical oxidant (electro-chlorination):

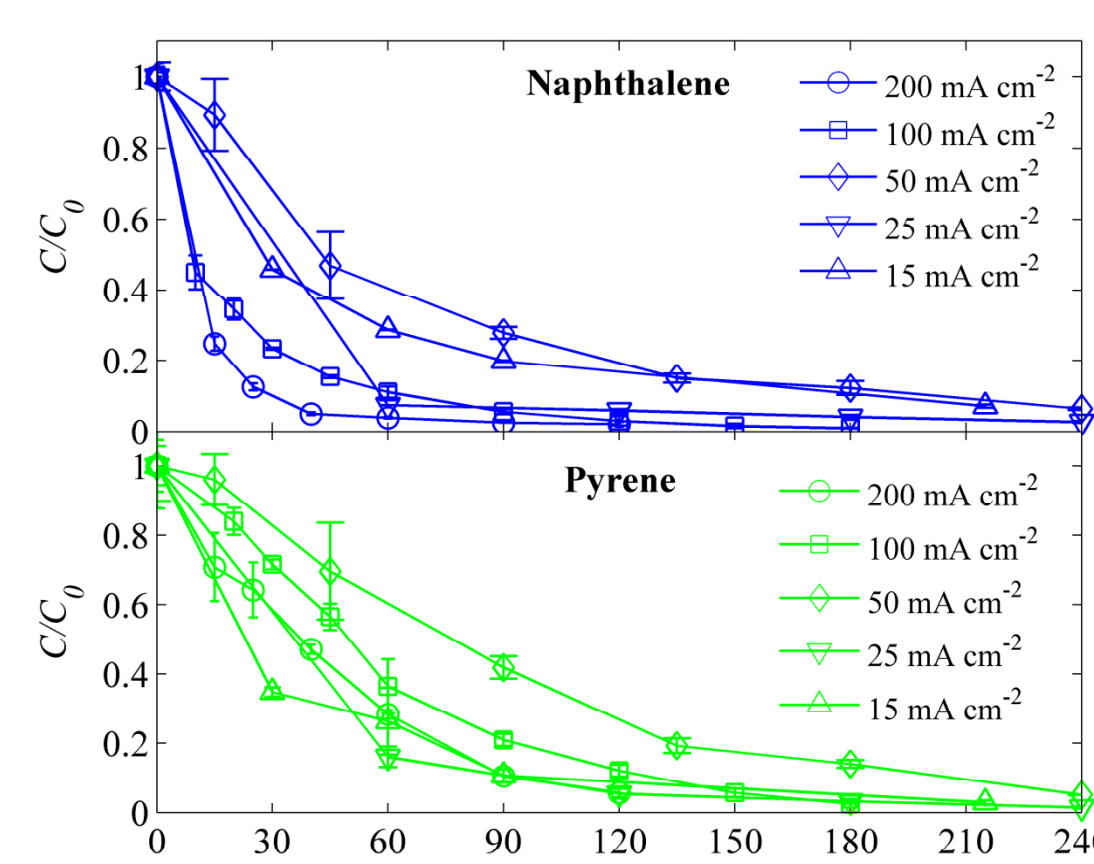
- Successful PAH removal obtained
- Energy consumption: 18.8 kWh m⁻³

Detailed model solution studies

Reaction kinetics & influence of current density



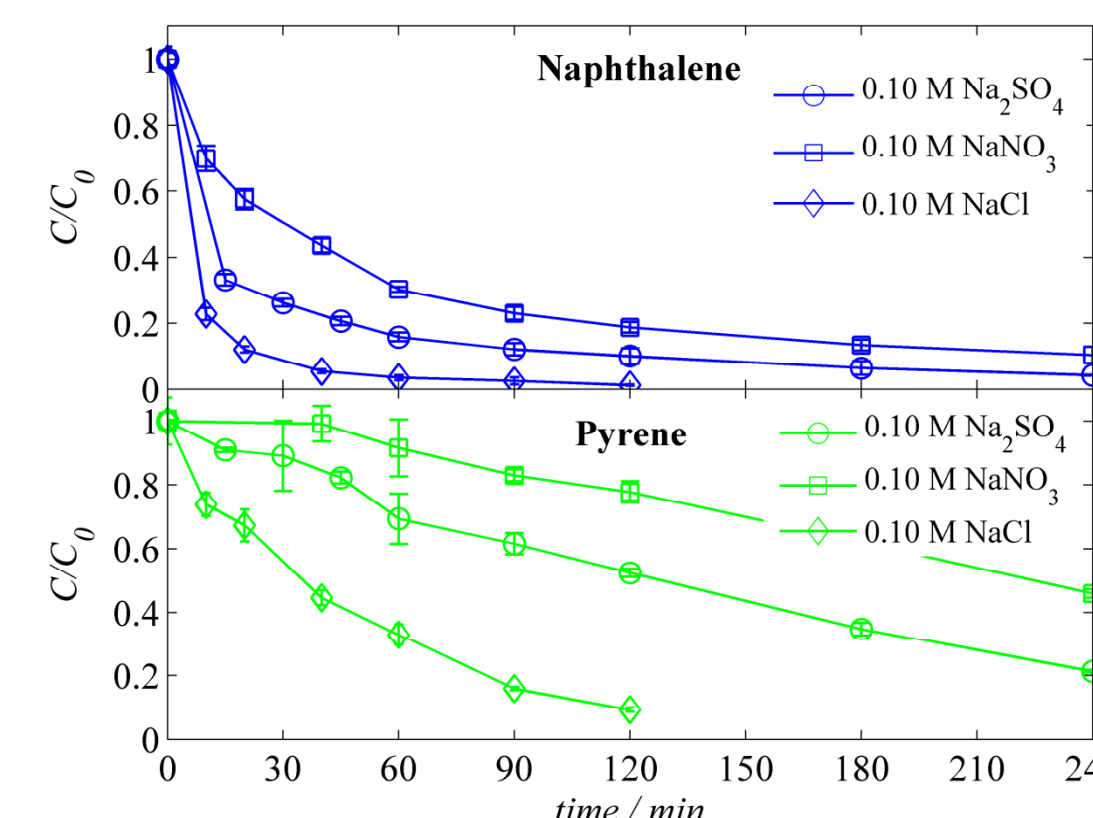
The evolution of the relative concentrations of (○) naphthalene, (□) fluoranthene, and (△) pyrene during electrochemical oxidation in 0.10 M Na₂SO₄ electrolyte at 200 mA cm⁻². Subplot: The standard kinetic analysis, which showed difference in reaction kinetics.



The influence of current density on the evolution in the relative concentrations during electrolysis in 0.14 M NaCl electrolyte.

Results & Discussion II

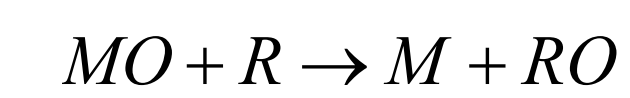
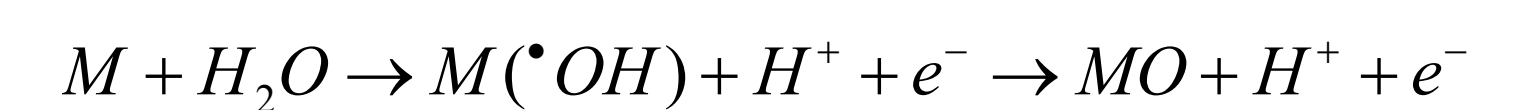
Influence of electrolyte



The influence of electrolyte composition on the evolution in the relative concentrations during electrolysis at a constant current density of 200 mA cm⁻². Due to a very similar behavior of fluoranthene and pyrene, only pyrene is shown in the plot.

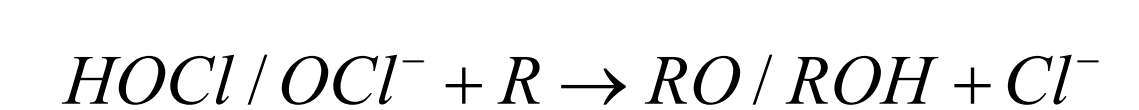
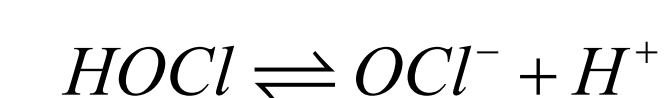
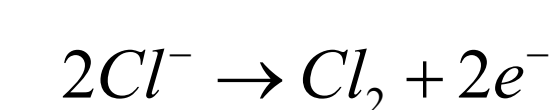
- Na₂SO₄ / NaNO₃ (Direct oxidation)

M: anode ; R: PAHs

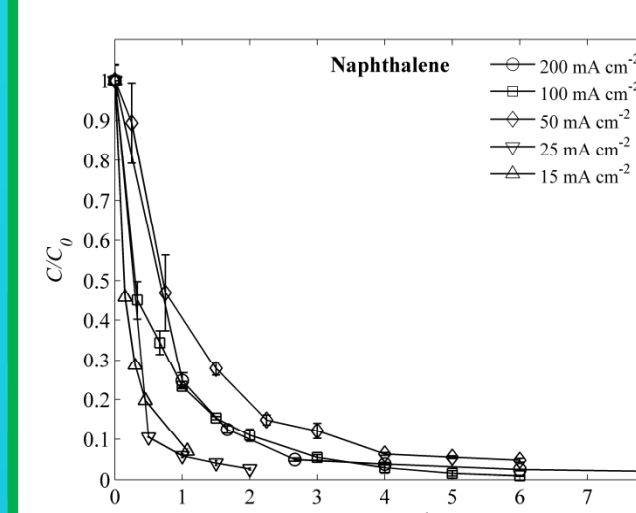


- NaCl (Indirect oxidation)

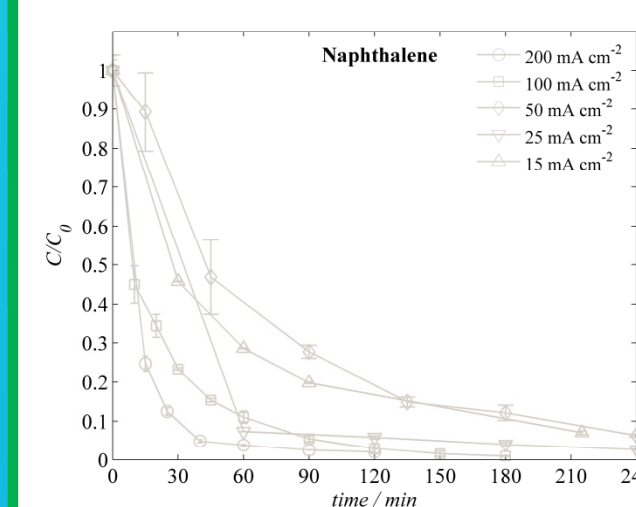
R: PAHs



Efficiency considerations



Degradation of naphthalene versus the amount of charge passed through the solution, Q (Ah L⁻¹).



Degradation of naphthalene versus time.

k_q: current efficiency constant

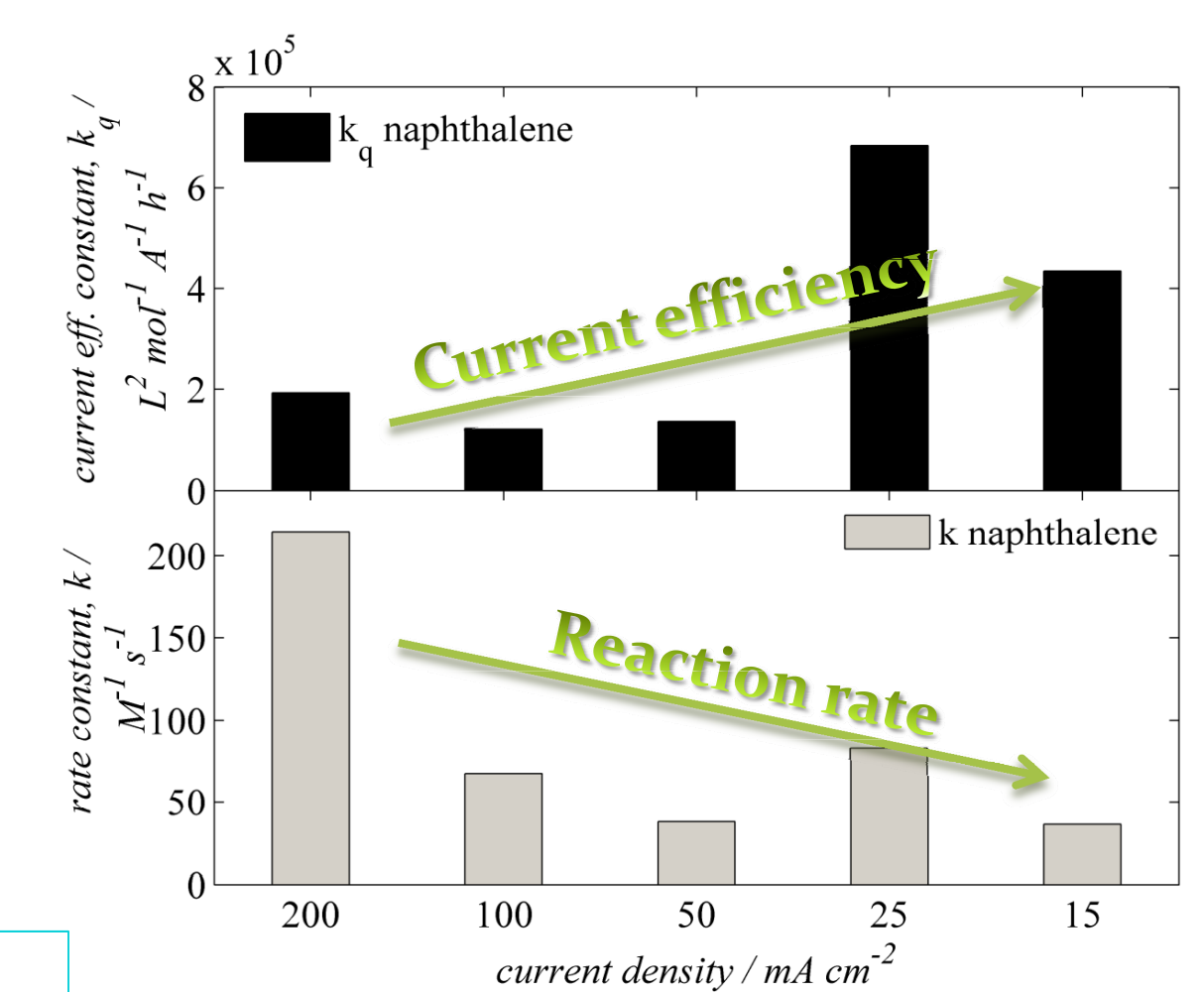
Relative measure of the efficiency of the oxidation of the specific PAH at different experimental settings

$$\frac{d[Naphthalene]}{dQ} = -k_q \cdot [Naphthalene]^2$$

k: kinetic rate constant

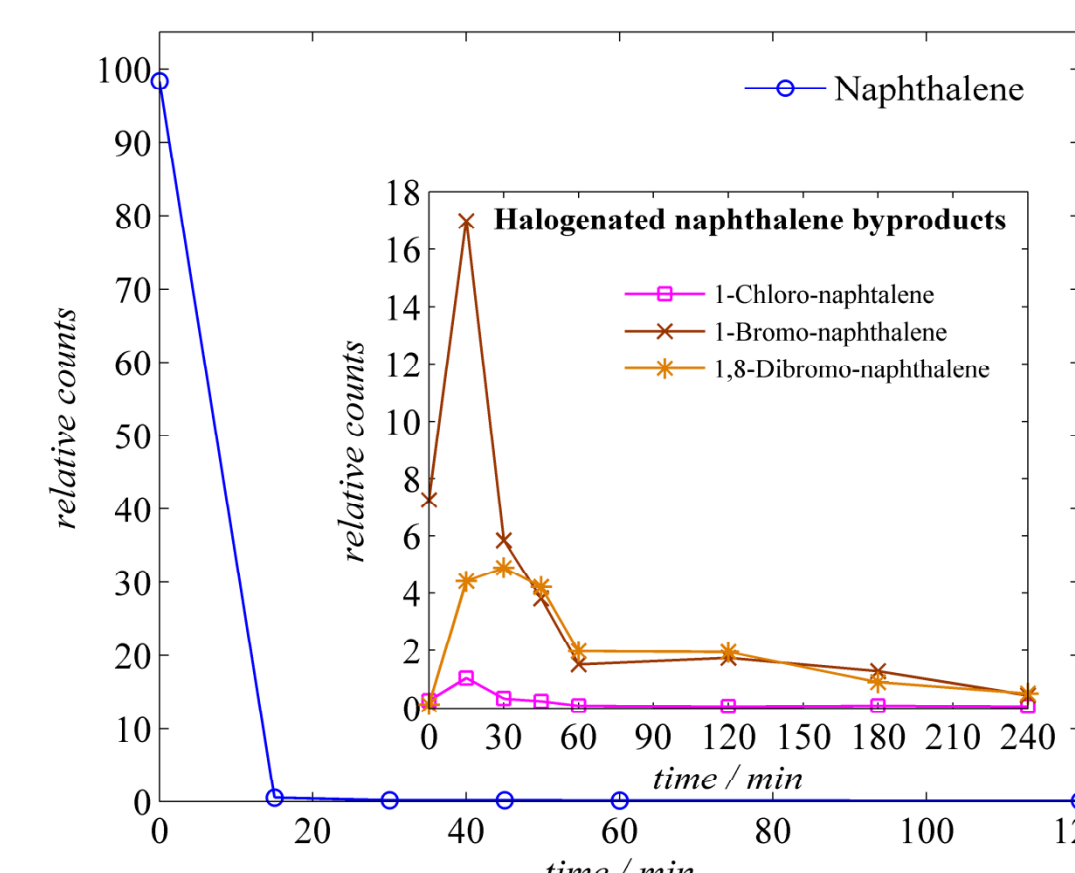
$$\frac{d[Naphthalene]}{dt} = -k \cdot [Naphthalene]^2$$

As the current density was decreased, the rate of reaction rate slowed down, but the current efficiency went up. At the low current densities, a higher yield of PAH degradation was obtained per spent ampere (lower cost), but the necessary reaction time was higher (larger reservoir dimensions).



Comparison of reaction rates and current efficiencies at different current densities for naphthalene in 0.14 M NaCl electrolytes. Naphthalene is chosen as an example, but the trends were similar for all three PAHs.

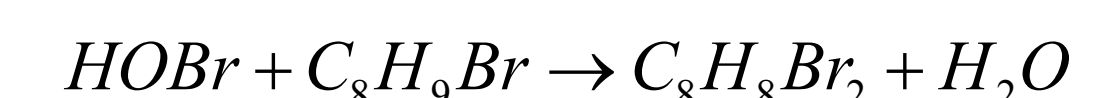
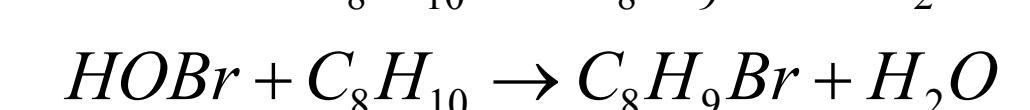
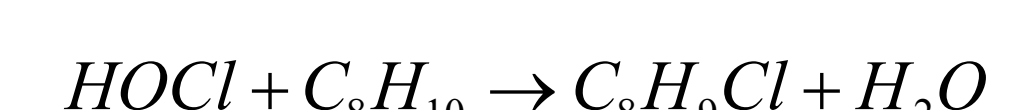
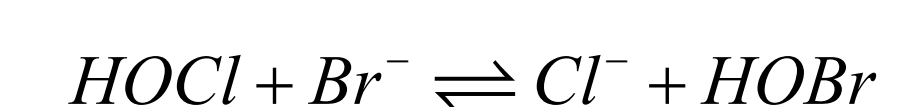
Halogenated byproduct formation



Removal of naphthalene and halogenated byproducts in 0.10 M NaCl + 0.001 M KBr (model sea water) at 25 mA cm⁻² and pH 6.

Formation of unwanted chlorinated and brominated byproducts is a risk when applying electrochemical oxidation in sea water. However, the low amount of mono- and dihalogenated naphthalene products detected during the treatment where all further degraded. Byproducts of fluoranthene and pyrene were not found.

- Possible routes of formation



Conclusions

Electrochemical purification of sediment run off water:

- Needed? In deed!!
- Feasible? Absolutely
- Expensive? Probably

Successful removal of PAHs was obtained by the electrochemical oxidation technique. Cost optimization is possible, which maybe will make the approach economically feasible. Halogenated byproducts are formed, but does not pose a significant risk due to subsequent degradation in the treatment process.

Acknowledgements



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